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Printing apparatus.

A transfer printing medium comprising a substrate supporting a thermal transfer dye and a radiation absorber positioned to provide thermal energy to the transfer dye when subjected to radiation within a predetermined absorption waveband, has a radiation absorber which is an infra-red absorbing poly(substituted)phthalocyanine compound in which each of at least five of the peripheral carbon atoms in the 1, 4, 5, 8, 9, 12, 13 or 16 positions (the «3,6-positions») of the phthalocyanine nucleus, as shown in Formula I, is linked by an atom from Group VB or Group VIB of the Periodic Table, other than oxygen, to a carbon atom of an organic radical. In preferred compounds each of the eight 3,6-positions is linked by an atom from Group VB or Group VIB, especially sulphur, selenium or nitrogan, to an organic radical.

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PRINTING APPARATUS

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The invention relates to laser transfer printing, and especially to apparatus suitable for printing multicolour designs and patterns.

Transfer printing is a technique which has been used for many years for printing patterns onto textiles and other receptor surfaces, and employs volatile or (more usually) sublimeable dyes, generally referred to collectively as "thermal transfer dyes". The thermal transfer dyes, usually in a formulation including a binder, are supported on a substrate such as paper, then, when eventually used, they are held firmly against the textile or other receptor surface and heat is applied to volatilise or sublime the dye onto that surface. The printing medium used for printing textiles thus usually comprises the various dyes printed onto the substrate in the form of the final pattern, and this is transferred by heating the whole area using a heated plate or roller. Thermal transfer dyes in a wide range of colours have been developed for such processes.

A more recent development is to use a laser as a source of energy for transferring the dyes. This enables just a single, very small, selected area to be heated at any one time, with only a corresponding small area of the dye being transferred, and by heating such selected areas in turn, the desired pattern can be built up, pixel by pixel, from a uniform sheet of printing medium. Computer control of such operations can enable complex designs of high definition to be printed at high speed, including multicolour designs by printing the different colours sequentially, either from different single colour sheets or from multicolour sheets carrying the different colours in different zones which can be brought into position in turn.

The transfer dyes can be heated directly by using a laser whose radiation lies within a strong absorption waveband of the dye, usually the complementary colour of the dye. However, this need to match the dye and the laser does restrict the choice of colours, and multicolour patterns require a corresponding number of lasers, one for each colour. The dyes can also be heated indirectly by incorporating a separate radiation absorber positioned to provide thermal energy to the transfer dyes when subjected to radiation within a predetermined absorption waveband, i.e. with writing radiation. This has previously been achieved by mixing carbon black with the transfer dye so that radiation of a wavelength different from that absorbed by the dye can be used. When printing with several colours, this has advantages in that the thermal energy produced is consistent with respect to the writing radiation irrespective of the colours used, and only a single laser is required. However we found that this did not prove entirely satisfactory because even though the carbon black would not sublime or volatilise like the dye, small particles did tend to be carried over with the dye molecules, thereby producing very obvious contamination.

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According to the present invention a transfer printing medium comprises a substrate supporting a thermal transfer dye and a radiation absorber positioned to provide thermal energy to the transfer dye when subjected to radiation within a predetermined absorption waveband, characterised in that the radiation absorber is a poly(substituted)phthalocyanine compound in which each of at least five of the

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peripheral carbon atoms in the 1,4, 5, 8, 9, 12, 13 or 16 positions of the phthalocyanine nucleus, as shown in Formula I is linked by an atom from Group VB or Group VIB of the Periodic Table, other than oxygen, to a carbon atom of an organic radical.

The specified poly(substituted)phthalocyanine compounds absorb in the near infra-red region of the electro-magnetic spectrum, e.g. from 750 to 1500 nm, but mainly from 750 to 1100 nm, with only very weak absorption in the visible region (i.e. within the range

of about 400-700 nm). The advantage of this is that should any of the present absorbers be carried over with the transfer dye during writing, it will not affect the colour balance of the transferred design. Moreover suitable infra-red lasers are available, including semiconductor diode lasers, which are generally cheap and can be matched to a range of dyes, and neodymium YAG lasers for giving radiation well into the near infra red at 1060 nm.

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The carbon atoms in the 1, 4, 5, 8, 9, 12, 13 and 16 positions are hereinafter referred to as the "3,6-carbon atoms" by relation to the equivalent 3,6-positions in the four molecules of phthalic anhydride, see Formula II, from which the phthalocyanine can be derived.

The remaining peripheral atoms of the phthalocyanine nucleus may be unsubstituted, i.e. carry hydrogen atoms, or be substituted by other groups, for example, halogen atoms or amino groups, or they may also be linked by an atom from Group VB or Group VIB of

the Periodic Table to a carbon atom of an organic radical. It is preferred that each of at least six, and more preferably at least eight, of the 3,6 carbon atoms is linked by a Group VB or Group VIB atom to an organic radical.

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The organic radical may be an optionally substituted aliphatic, alicyclic or aromatic radical and is preferably an optionally substituted aromatic radical, especially from the benzene, naphthalene and mono- or bi-cyclic, heteroaromatic series. Examples of suitable aromatic radicals are optionally substituted phenyl, phenylene, naphthyl, especially naphth-2-yl, naphthylene, pyridyl, thiophenyl, furyl, pyrimidyl and benzthiazolyl. Aliphatic radicals are preferably from the alkyl and alkenyl series containing up to 20 carbon atoms, such as vinyl, allyl, butyl, nonyl, dodecyl, octadecyl and octadecenyl. Alicyclic radicals are preferably homocyclic containing from 4 to 8 carbon atoms, such as cyclohexyl. The organic radical may be monovalent and attached to a single peripheral carbon atom through a single Group VB or Group VIB atom or it may be polyvalent, preferably divalent, and attached to adjacent peripheral carbon atoms through identical or different atoms from Group VB and Group VIB. Where the organic radical is polyvalent it may be attached to two or more phthalocyanine nuclei.

Examples of substituents for the aromatic and heteroaromatic radicals are alkyl, alkenyl, alkoxy and alkylthio, and halo substituted derivatives thereof, especially those containing up to 20 carbon atoms, aryl, arylthio, especially phenyl and phenylthio, halogen, nitro, cyano, carboxyl, aralkyl, aryl- or

alkyl-sulphonamido, aryl- or alkyl-sulphone, aryl- or alkyl-sulphoxide, hydroxy and primary, secondary or tertiary amino. Examples of substituents for the aliphatic and cycloaliphatic radicals are alkoxy, alkylthio, halo, cyano and aryl. In these substituents the alkyl and alkenyl groups preferably contain up to 20, and more preferably up to 4, carbon atoms and the aryl groups are preferably mono- or bi-homo- or hetero-cyclic. Specific examples of substituents are methyl, ethyl, dodecyl, methoxy, ethoxy, methylthio, allyl, trifluoromethyl, bromo, chloro, fluoro, benzyl, COOH, -COOCH3, -COOCH2C6H5, -NHSO2CH3, -SO2C6H5, NH2, -NHC2H5, and H(CH3)2.

Examples of suitable atoms from Group VB and Group VIB for linking the organic radical to a peripheral carbon atom of the phthalocyanine nucleus are sulphur, selenium, tellurium and nitrogen or any combination of these. Where an organic radical is linked to adjacent peripheral carbon atoms the second bridging atom may be any atom from Group VB or Group VIB and examples are sulphur, oxygen, selenium, tellurium and nitrogen. Where the linking atom is nitrogen the free valency may be substituted or unsubstituted, e.g. it may carry an alkyl group, preferably C_{1-4} -alkyl or an aryl group, preferably phenyl.

The phthalocyanine compounds of the present invention can be prepared by heating a phthalocyanine compound carrying halogen atoms attached to the peripheral carbon atoms to which it is wished to attach the Group VB or Group VIB atoms, with at least six equivalents of an organic thiol or an equivalent

compound in which the sulphur in the thiol group is replaced by selenium (selenol), tellurium (tellurol) or NT (amine), in an organic solvent.

The organic solvent, which need not necessarily be a liquid at ambient temperatures and may only partially dissolve the reactants, preferably has a boiling point from 100°C to 300°C and more preferably from 150°C to 250°C. The organic solvent is preferably essentially inert although it may catalyse the reaction. Examples of suitable solvents are methylcyclohexanol, octanol, ethylene glycol, and especially benzyl alcohol and quinoline.

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Reaction is conveniently carried out under reflux, preferably from 100°C to 250°C and more preferably above 150°C, in the presence of an acid binding agent, such as potassium or sodium hydroxide or sodium carbonate, to neutralise the halo acid formed. The product may be isolated by filtration or by distillation of the organic liquid. The isolated product is preferably purified by repeated recrystallisation from a suitable solvent, such as ethanol, chloroform or pyridine, and/or chromatography, using a silica-filled column and an aromatic solvent, such as toluene or xylene, as eluent.

The phthalocyanine nucleus may be metal free, i.e. it may carry two hydrogen atoms at the centre of the nucleus, or it may be complexed with a metal or oxy-metal derivative, i.e. it may carry one or two metal atoms or oxy-metal groups complexed within the centre of the nucleus. Examples of suitable metals and oxy-metals are copper, lead, cobalt, nickel, iron, zinc, germanium, indium, magnesium, calcium, palladium, gallium and vanadium.

The radiation absorber and transfer dye are preferably intimately mixed in a common coating layer on the supporting substrate. However, an alternative arrangement that can also work is one in which they are arranged as separate layers on the same side of the substrate, preferably with the radiation absorber forming the layer nearer to the substrate.

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For supporting the dyes in the printing medium we prefer to use a polyester film, such as Melinex film, to take advantage of its high transparency in the near infra-red, and its generally good heat stability. EXAMPLES

The following poly(substituted)phthalocyanine compounds were prepared and their absorption maxima measured as solutions in chloroform (Chlor), toluene (Tol) or after deposition on glass (Glass) unless otherwise indicated. Extinction coefficients were determined in toluene or the only solvent in which the absorption maximum was recorded.

20	Example	e Product	Absorption				
			Maxima (nm)			Extinction	
			Chlor	Tol	Glass	Coefficient	
	1	octa-3,6-(4-methyl-					
		phenylthio)-H ₂ Pc	813	805	828	170,000	
25	2	octa-3,6-(4-methyl-			-		
		thio)-CuPc	797	787	797	156,000	
	3	octa-3,6(3-methyl-					
		phenylthio)H ₂ Pc	805	797	818	160,000	
	4	hepta-3,6(4-t-butyl-					
30		phenylthio)H ₂ Pc	798	790		173,000	
	5	octa-3,6(4-t-butyl-					
		phenylthio)H ₂ Pc	793		797	152,000	

	Exampl	e Product	Absorption				
			Maxima (nm)			Extinction	
			Chlor	Tol	Glass	Coefficient	
	6	octa-3,6(4-t-buty1-					
		phenylthio)CuPc	803		797	216,000	
	7	hepta-3,6(4-n-nonyl-					
		phenylthio)H ₂ Pc	800		809		
5	8	hepta-3,6(4-dodecy1-					
		phenylthio)H ₂ Pc	789	787	795		
	9	hexa-3,6(3,4-dimethyl-					
		phenylthio)H ₂ Pc	807	803	830		
	10	octa-3,6(4-methoxy-					
10		phenylthio)H ₂ Pc	799	792		161,500	
	11	octa-3,6(4-methoxy-					
		phenylthio)CuPc	805		813	155,000	
	12	octa-3,6(4-butoxy-					
		phenylthio)CuPc	800	786			
15	13	octa-3,6(4-dodecyloxy-					
		phenylthio)H ₂ Pc	818	808	859		
	14	octa-3,6(4-dodecyloxy-					
		phenylthio)CuPc	807	794	822		
	15	octa-3,6(naphth-2-					
20		ylthio)CuPc	799		796	136,000	
	16	octa-3,6(4-octoxy-					
	•	phenylthio)H ₂ Pc	816	806	846		
	17	penta-3,6(4-octoxy-					
		phenylthio)CuPc	775				
25	18	pentadeca(4-methy1-					
		thio)-CuPc	775	768	790	169,000	
	19	deca(4-methylthio)-					
		pentachloro-CuPc	758	752	770	174,000	
	20	pentadeca(t-butyl-					
30		phenylthio)CuPc	774	760	784	142,000	
	21	pentadeca(3-methy1-					
		phenylthio)CuPc	771	766	786		

	Exampl	e Product	Absorption			
			Maxima (nm)		n)	Extinction
			Chlor	Tol	Glass	Coefficier
	22	pentadeca(4-methoxy-				
		phenylthio)CuPc	786		801	190,000
	23	terdeca(4-butoxy-				
		phenylthio)CuPc	775	768	797	158,000
5	24	pentadeca(4-butoxy-				
		phenythio)CuPc	786	780	801	182,000
	25	pentadeca(4-dodecoxy-				
		phenylthio)CuPc	778	770	792	162,000
	26	pentadeca(phenylthio)				
10		CuPc	772	768	794	
	27	tetradeca(2-methoxy-				
		phenylthio)CuPc	770			
	28	pentadeca(4-methyl-				
		thiophenylthio)CuPc	788	784	810	208,500
15	29	deca(4-ethylthio-	\			
		phenylthio)CuPc	756	752		
	30	pentadeca(4-chloro-				
	•	phenylthio)CuPc	774		787	181,000
	31	unadeca(4-dimethyl-				
20		aminophenylthio)CuPc	782		805	118,000
	32	terdeca(naphth-1-				
		ylthio)CuPc	765	760		
	33	pentadeca(naphth-2-				
		ylthio)CuPc	786	781	799	197,000
25	34	pentadeca(phenyl-				
		seleno)CuPc	776			
	35	hexadeca(4-methyl-				
		phenyl-thio)PbPc	769		792	
	36	hexadeca(4-methyl-				
30		phenylthio)H ₂ Pc	769			
	37	hexadeca(4-methyl-				
		phenylthio)CuPc	778	770	796	220,000

	Exampl	e Product	Absorption				
			Maxim	a (nm	Extinction		
			Chlor	Tol	Glass	Coefficient	
	38	hexadeca(4-methyl-					
		phenylthio)ZnPc	768		791		
	39	hexadeca(4-chloro-					
		phenylthio)CuPc	770		789	220,000	
5	40	deca(naphth-2-ylthio)					
		H ₂ Pc	744				
	41	hepta(4-methylphen-1,					
		2-ylene-dithio)-di(4-					
		methyl-2-thiolphenyl-					
10		thio)-H ₂ Pc	800	797	832	94,000	
	42	hepta(4-methylphen-1,					
		2-dithio-ylene)-mono					
		(4-methyl-2-thio-					
		phenylthio)-CuPc	790	787	828	91,000	
15	43	penta(phen-l-amino-2-					
		thio-ylene)-penta(2-				•	
		aminophenylthio)-CuPc	909	(in p	yridi	ne)	
	44	pentadeca(ethylthio)-					
		monoisoamyloxy-H2Pc	804	807	827		
20	45	hexadeca(cyclohexyl-					
		thio)-ZnPc	846	852	860	95,000	
	46	tetradeca(ethylthio)					
		monoamyloxy-H ₂ Pc	801	802			
	47	(ethylthio) _{15.3}					
25		(amyloxy)0.7-H2Pc	805	808	830	149,000	
	48	hexadeca(n-propyl-					
		thio)-H ₂ Pc	802	800	819	157,600	
	49	pentadeca(i-propyl-					
		thio)monoamyloxy-H ₂ Pc	809		823	136,500	
30	50	pentadeca(n-butyl-					
		thio)monoamyloxy-H ₂ Pc	807		817	147,000	

	Examp	Le	Product	Absorption				
				Maxima (nm)		<u>a)</u>	Extinctio	
				Chlor	Tol	Glass	Coefficie	
	51	pentadeo	ca(n-pentyl-					
		thio)mor	noamyloxy-H ₂ Pc	802	802		162,500	
	52	octa(but	ylthio)octa					
		(ethylth	nio)-H ₂ Pc	809	805	815	129,000	
5	53	octa(but	ylthio)octa					
		(ethylth	nio)-H ₂ Pc	803	797	815	115,500	
	54	pentadeo	a(cyclohexyl-					
		thio)mor	oamyloxy-H ₂ Pc	812	810	818	120,000	
	55	hexadeca	(n-octylthio)					
10		-H2Pc		818	811	•		
	56	pentadec	a(s-butyl-					
		thio)mon	oamyloxy-H ₂ Pc	805	801	-	133,000	
	57	pentadeo	a(benzylthio)					
		monoamyl	oxy-H ₂ Pc	810	809		84,000	
15	58	hexadeca	(phenylthio)					
		-H ₂ Pc		790				
	59	octa-3,6	-(isopropyl-					
		thio)-H ₂	Pc	802			167,000	
	60	pentadec	a(n-propyl-					
20		thio)mon	oamyloxy-CuPc	783	785	805	170,500	
	61	pentadec	a(n-pentyl-					
		thio)mon	oamyloxy-CuPc	784	783		182,000	
	62	pentadec	a(cyclohexyl-					
		thio)mon	oamyloxy-CuPc	789	781	803	163,000	
25	63	pentadec	a-s-butyl-					
		thio)mon	oaryloxy-CuPc	787	778		168,000	
	64	pentadec	a(benzylthio)					
		monoaryl	oxy-CuPc	797	789		109,000	
	65	pentadec	a(cyclohexyl-					
30		thio)mon	oamyloxy-PbPc	838	830	840	111,000	
	66	octapipe	ridino-octa-					
		chloro-H	2 ^{PC}	835				

CLAIMS

a substrate supporting a thermal transfer dye and a radiation absorber positioned to provide thermal energy to the transfer dye when subjected to radiation within a predetermined absorption waveband, characterised in that the radiation absorber is a poly(substituted)phthalocyanine compound in which each of at least five of the peripheral carbon atoms in the 1, 4, 5, 8, 9, 12, 13 or 16 positions of the phthalocyanine nucleus, as shown in Formula I is linked by an atom from Group VB or Group VIB of the Periodic Table, other than oxygen, to a carbon atom of an organic radical.

A transfer printing medium as claimed in Claim

1, characterised in that each of the eight peripheral
carbon atoms in the 1, 4, 5, 8, 9, 12, 13 and 16
positions of the phthalocyanine nucleus is linked by an
atom from Group VB or Group VIB of the Period Table,
other than oxygen, to a carbon atom of an organic
radical.

- 3 A transfer printing medium as claimed in Claim 2 wherein the remaining peripheral carbon atoms of the phthalocyanine nucleus are unsubstituted.
- A transfer printing medium as claimed in any one of Claims 1 to 3 wherein the organic radical is an optionally substituted aliphatic, cycloaliphatic or aromatic radical.
- 5 A transfer printing medium as claimed in Claim 4 wherein the organic radical is an optionally substituted phenyl, naphthyl or mono- or bi-cyclic heteroaromatic radical.
- A transfer printing medium as claimed in any one of Claims 1 to 5 wherein the organic radical is bivalent and attached to adjacent peripheral carbon atoms on the phthalocyanine nucleus through atoms from Group VB or Group VIB of the Periodic Table.
- 7 A transfer printing medium as claimed in any one of Claims I to 6 wherein the atom from Group VB or Group VIB of the Periodic Table is sulphur, selenium, tellurium or nitrogen.
- A transfer printing medium as claimed in any one of Claims 1 to 7 wherein the radiation absorber and thermal transfer dye are intimately mixed in a common coating layer on the supporting substrate.
- 9 A transfer printing medium as claimed in any one of Claims 1 to 8 wherein the substrate is a polyester film transparent to radiation in the near infra red.